WHAT IS CLAIMED IS:

- 1. A transgenic cotton plant or seed, cells or tissues thereof comprising event EE-GH1 in its genome.
- 2. The transgenic cotton plant, or seed, cells or tissues of claim 1, the genomic DNA of which, when analyzed using the Elite event identification protocol for EE-GH1 with two primers comprising the nucleotide sequence of SEQ ID NO: 2 and SEQ ID NO: 3 respectively, yields a DNA fragment of between 250 and 290 bp.
- 3. The cotton plant, or seed, cells or tissues thereof, according to claim 2, wherein said DNA fragment is a fragment of about 269 bp.
- A cotton plant, or seed, cells or tissues thereof, obtained by propagation of and/or breeding with a cotton plant grown from the seed deposited at the ATCC under accession number PTA-3343.
- 5. A cotton plant, seed, cells or tissues thereof which is the progeny of the seed deposited at the ATCC under accession number PTA-3343
- 6. A method for identifying elite event EE-GH1 in biological samples, which method comprises detecting an EE-GH1 specific region with a primer or probe which specifically recognizes the 5' flanking region of SEQ ID NO: 3 or the 3' flanking region of SEQ ID NO: 4 of EE-GH1.
- 7. The method of claim 6, said method comprising amplifying a DNA fragment of between 100 and 350 bp from a nucleic acid present in said biological samples using a polymerase chain reaction with at least two primers, one of which recognizes the 5' or 3' flanking region of EE-GH1 and the other which recognizes a sequence within the foreign DNA of EE-GH1.

- 8. The method of claim 7, wherein said one primer recognizes a sequence within the 5' flanking region of SEQ ID NO: 3 and said other primer recognizes a sequence within the foreign DNA of EE-GH1.
- 9. The method of claim 8, wherein said primer recognizing a sequence within the 5' flanking region of EE-GH1 comprises the sequence of SEQ ID NO: 2.
- 10. The method of any one of claims 6 to 9, wherein said primer recognizing a sequence within the foreign DNA comprises the sequence of SEQ ID NO: 1.
- 11. A method for identifying EE-GH1 in a biological sample, which method comprises detecting an EE-GH1 specific region with a specific primer or probe which hybridizes under stringent conditions to a sequence within the 5' of SEQ ID NO: 3 or within the 3' flanking sequence of SEQ ID NO: 4 of EE-GH1.
- 12. A method for identifying a transgenic plant, or cells or tissues thereof, comprising the elite event EE-GH1, which method comprises establishing that genomic DNA can be used, according to a PCR identification protocol, to amplify a DNA fragment of between 250 and 290 bp, using a polymerase chain reaction with two primers having the nucleotide sequence of SEQ ID NO: 1 and SEQ ID NO: 2, respectively.
- 13. A kit for identifying elite event EE-GH1 in biological samples, said kit comprising at least one PCR primer or probe, which recognizes a sequence within the 5' flanking region of SEQ ID NO: 3 or the 3' flanking region of SEQ ID NO: 4 of EE-GH1.
- 14. The kit of claim 13, wherein said at least one PCR primer recognizes a sequence within the plant DNA in SEQ ID NO: 3.

- 15. The kit of claim 14, wherein said primer recognizing a sequence within the plant DNA in SEQ ID NO: 3 comprises the sequence of SEQ ID NO: 2.
- 16. The kit of Claims 13 to 15, which further comprises at least a second PCR primer or probe which recognizes a sequence within the foreign DNA of EE-GH1.
- 17. The kit of claim 16, wherein said primer recognizing a sequence within the foreign DNA of EE-GH1 comprises the sequence of SEQ ID NO: 1.
- 18. A method for confirming seed purity, which method comprises detecting an EE-GH1 specific DNA sequence with a specific primer or probe which specifically recognizes a sequence within the 5' flanking region of SEQ ID NO: 3 or the 3' flanking region of SEQ ID NO: 4 of EE-GH1, in seed samples.
- 19. A method for screening seeds for the presence of EE-GH1, which method comprises detecting an EE-GH1 specific DNA sequence with a specific primer or probe which specifically recognizes a sequence within the 5' flanking region of SEQ ID NO: 3 or the 3' flanking region of SEQ ID NO: 4 of EE-GH1, in samples of seed lots.
- 20. A seed deposited at the ATCC under accession number PTA-3343.
- 21. A cotton seed comprising elite event EE-GH1, reference seed comprising said event having been deposited at the ATCC under accession number PTA-3343.
- 22. A cotton plant, cell or tissue or plant material thereof comprising elite event EE-GH1, derived from the seed of claim 21.

- 23. Transgenic cotton plants, seeds, cells or tissues, the genomic DNA of which comprises a transgene integrated into the chromosomal DNA in a region which comprises a sequence of at least 40 bp which hybridizes under stringent conditions with a sequence which is complementary to the sequence of SEQ ID NO: 5.
- 24. A process for producing a transgenic cotton plant or cell or tissue of a cotton plant, said process comprising introducing a recombinant DNA molecule into a region of cotton chromosomal DNA corresponding to a sequence of at least 40 bp that hybridizes under stringent conditions with a sequence that is complementary to the sequence of SEQ ID NO: 5, and, optionally, regenerating a cotton plant from the transformed cotton cell or tissue.
- 25. The process of claim 24, wherein said recombinant DNA molecule comprises an herbicide resistance gene.
- 26. The plant or cell or tissue of a cotton plant obtained by the process of claims 24 or 25.
- 27. A transgenic cotton plant or seed, cells or tissues thereof comprising
 - (i) event EE-GH1 in its genome ;or
- (ii) event EE-GH1 with the proviso that the *bar* gene used in the event is substituted with a nucleic acid sequence that hybridizes to the complement of the *bar* gene under stringent conditions.